

**A METHOD FOR PRODUCING, IN YEAST, A HYDROXYLATED  
TRIPLE HELICAL PROTEIN, AND YEAST HOST CELLS USEFUL  
IN SAID METHOD**

5 This is a Continuation-in-part Application of Application No. 09/297,269, filed  
28 April, 1999, which in turn is a National stage filing under 35 U.S.C. § 371 of  
PCT/AU97/00721, filed October 29, 1997.

**Field of the Invention:**

10 This invention relates to the production of hydroxylated triple helical proteins  
such as natural and synthetic collagens, natural and synthetic collagen fragments, and  
natural and synthetic collagen-like proteins, by recombinant DNA technology. In  
particular, the invention relates to a method for producing hydroxylated triple helical  
15 proteins in yeast host cells by introducing to a suitable yeast host cell, DNA sequences  
encoding the triple helical protein as well as prolyl 4-hydroxylase (P4H), in a manner  
wherein the introduced DNA sequences are replicated, stably retained and segregated  
by the yeast host cells.

**Background of the Invention:**

20 The collagen family of proteins represents the most abundant protein in  
mammals, forming the major fibrous component of, for example, skin, bone, tendon,  
cartilage and blood vessels. Each collagen protein consists of three polypeptide chains  
(alpha chains) characterised by a (GlyXY)<sub>n</sub> repeating sequence, which are folded into a  
triple helical protein conformation. Type I collagen (typically found in skin, tendon,  
25 bone and cornea) consists of two types of polypeptide chain termed  $\alpha 1(I)$  and  $\alpha 2(I)$   
[i.e.  $\alpha 1(I)_2\alpha 2(I)$ ], while other collagen types such as Type II [ $\alpha 1(II)_3$ ] and Type III  
[ $\alpha 1(III)_3$ ] have three identical polypeptide chains. These collagen proteins  
spontaneously aggregate to form fibrils which are incorporated into the extracellular  
matrix where, in mature tissue, they have a structural role and, in developing tissue,  
30 they have a directive role. The collagen fibrils, after cross-linking, are highly insoluble  
and have great tensile strength.

The ability of collagen to form insoluble fibrils makes them attractive for  
numerous medical applications including bioimplant production, soft tissue  
augmentation and wound/burn dressings. To date, most collagens approved for these  
35 applications have been sourced from animal sources, primarily bovine. While such  
animal-sourced collagens have been successful, there is some concern that their use

risks serious immunogenicity problems and transmission of infective diseases and spongiform encephalopathies (e.g. bovine spongiform encephalopathy (BSE)). Accordingly, there is significant interest in the development of methods of production of collagens or collagen fragments by recombinant DNA technology. Further, the use of recombinant DNA technology is desirable in that it allows for the potential production of synthetic collagens and collagen fragments which may include, for example, exogenous biologically active domains (i.e. to provide additional protein function) and other useful characteristics (e.g. improved biocompatibility and stability).

The *in vivo* biosynthesis of collagen proteins is a complex process involving many post translational events. A key event is the hydroxylation by the enzyme prolyl 4-hydroxylase (P4H) of prolyl residues in the Y-position of the repeating (GlyXY)<sub>n</sub> sequences to 4-hydroxyproline. This hydroxylation has been found to be beneficial for nucleation of folding of triple helical proteins. For collagens, it is essential for stability at body temperature. Accordingly, the development of a commercially viable method for the production of recombinant collagen requires co-expression of P4H with the alpha chains. For mammalian host cells, co-expression of P4H will occur autonomously since these cells should naturally express P4H. However, for yeast host cells, which for reasons of cost, ease and efficiency are more attractive for expression of recombinant eukaryotic proteins, transformation with DNA sequences encoding P4H will also be required. Since P4H consists of  $\alpha$  and  $\beta$  subunits of about 60 kDa and 60 kDa, yeast host cells for expression of recombinant collagen will require co-transformation with at least three exogenous DNA sequences (i.e., encoding an alpha chain, P4H  $\alpha$  subunit and P4H  $\beta$  subunit) and stability problems would therefore be expected if cloned on three separate vectors or, alternatively, all on episomal type vector. Indeed, even under continuous selection pressure, many episomal type vectors suffer stability problems if they are large or are present at relatively low copy number. An object of the present invention is therefore to provide a method for expressing recombinant collagen and other triple helical proteins from yeast host cells wherein the introduced DNA sequences are stably retained and segregated independent of continuous selection pressure.

#### **Summary of the Invention:**

Thus, in a first aspect, the present invention provides a method of producing, in yeast, a hydroxylated triple helical protein, said method comprising the steps of:

(A) introducing into a suitable yeast host cell:

(i) a first DNA molecule comprising a DNA sequence encoding prolyl 4-hydroxylase I-subunit (P4HI) operably linked to a promoter functional in said yeast host cell,

(ii) a second DNA molecule comprising a DNA sequence encoding prolyl 4-hydroxylase 9-subunit (P4H9) operably linked to a promoter functional in said yeast host cell, and

(iii) a third DNA molecule comprising a DNA sequence encoding a polypeptide or peptide operably linked to a promoter functional in said yeast host cell, wherein said polypeptide or peptide is one which, when hydroxylated, forms said hydroxylated triple helical protein, and wherein said polypeptide or peptide is a synthetic polypeptide or peptide represented by the following formula:

$$(A)_l - (B)_m - [Z] - (C)_o - (D)_p,$$

wherein;

Z is a domain comprising two or more repeat units of the formula:

$$[(E)_q - (GlyXY)_i - (F)_r],$$

wherein;

E and F represent sequences of one or more amino acids, which sequences may vary from repeat unit to repeat unit, and for each repeat unit q and r are each independently selected from 0 and 1, and

i is  $\geq 1$  such that domain Z comprises 2 to 1500 GlyXY triplets,

Gly represents glycine, and

X and Y, which may be the same or different, represent an amino acid, and wherein the identity of each amino acid represented by X and Y may vary from GlyXY triplet to GlyXY triplet, but wherein at least one Y of the  $(GlyXY)_i$  sequence must be proline,

A and D, which may be the same or different, each represent a polypeptide or peptide domain which optionally comprises a triple helical forming repeating sequence  $(GlyXY)_n$ , and l and p are each independently selected from 0 and 1,

B and C, which may be the same or different, each represent a polypeptide or peptide domain which is heterologous to collagen proteins and which does not comprise a triple helical forming repeating sequence  $(GlyXY)_n$ , and m and o are each independently selected from 0 and 1; and

(B) culturing the resulting yeast host cell of step (A) under conditions suitable to express said P4HI and P4H9 and said synthetic polypeptide or peptide, to produce said hydroxylated triple helical protein;

wherein during culturing in step (B), each of said first DNA molecule, said second DNA molecule and said third DNA molecule are replicated, stably retained and segregated by the yeast host cell.

In a second aspect, the present invention provides a yeast host cell capable of producing a hydroxylated triple helical protein, said yeast host cell including:

(i) a first DNA sequence encoding prolyl 4-hydroxylase I-subunit (P4HI) operably linked to a promoter functional in said yeast host cell,

(ii) a second DNA sequence encoding prolyl 4-hydroxylase 9-subunit (P4H9) operably linked to a promoter functional in said yeast host cell, and

(iii) a third DNA sequence encoding a polypeptide or peptide operably linked to a promoter functional in said yeast host cell, wherein said polypeptide or peptide is one which, when hydroxylated, forms said hydroxylated triple helical protein, and wherein said polypeptide or peptide is a synthetic polypeptide or peptide represented by the following formula:

$$(A)_1 - (B)_m - [Z] - (C)_o - (D)_p,$$

wherein;

Z is a domain comprising two or more repeat units of the formula:

$$[(E)_q - (GlyXY)_i - (F)_r],$$

wherein;

E and F represent sequences of one or more amino acids, which sequences may vary from repeat unit to repeat unit, and for each repeat unit q and r are each independently selected from 0 and 1, and

i is  $\geq 1$  such that domain Z comprises 2 to 1500 GlyXY triplets,

Gly represents glycine, and

X and Y, which may be the same or different, represent an amino acid, and wherein the identity of each amino acid represented by X and Y may vary from GlyXY triplet to GlyXY triplet, but wherein at least one Y of the (GlyXY)<sub>i</sub> sequence must be proline,

A and D, which may be the same or different, each represent a polypeptide or peptide domain which optionally comprises a triple helical forming repeating sequence (GlyXY)<sub>n</sub>, and l and p are each independently selected from 0 and 1,

5 B and C, which may be the same or different, each represent a polypeptide or peptide domain which is heterologous to collagen proteins and which does not comprise a triple helical forming repeating sequence (GlyXY)<sub>n</sub>, and m and o are each independently selected from 0 and 1; and

10 wherein each of said first DNA sequence, said second DNA sequence and said third DNA sequence are replicated, stably retained and segregated by the yeast host cell.

In a third aspect, the present invention provides an hydroxylated triple helical protein comprising a polypeptide or peptide which is a synthetic polypeptide or peptide represented by the following formula:

$$15 \quad (A)_l - (B)_m - [Z] - (C)_o - (D)_p,$$

wherein;

Z is a domain comprising two or more repeat units of the formula:

$$20 \quad [(E)_q - (GlyXY)_i - (F)_r],$$

wherein;

25 E and F represent sequences of one or more amino acids, which sequences may vary from repeat unit to repeat unit, and for each repeat unit q and r are each independently selected from 0 and 1, and

i is ≥ 1 such that domain Z comprises 2 to 1500 GlyXY triplets,

Gly represents glycine, and

30 X and Y, which may be the same or different, represent an amino acid, and wherein the identity of each amino acid represented by X and Y may vary from GlyXY triplet to GlyXY triplet, but wherein at least one Y of the (GlyXY)<sub>i</sub> sequence must be proline,

A and D, which may be the same or different, each represent a polypeptide or peptide domain which optionally comprises a triple helical forming repeating sequence (GlyXY)<sub>n</sub>, and l and p are each independently selected from 0 and 1,

35 B and C, which may be the same or different, each represent a polypeptide or peptide domain which is heterologous to collagen proteins and which does not

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**Detailed disclosure of the Invention:**

Preferably, the method of the invention is achieved by including the exogenous nucleotide sequences within a vector(s) including a CEN sequence. Particularly preferred are the CEN sequence-including YAC (yeast artificial chromosome) vectors (Cohen *et al.*, 1993) and pYEUra3 vectors (Clontech, Cat. No 6195-1). Other vectors including a CEN sequence may be generated by cloning a CEN sequence into any suitable expression vector.

Where one or more of the exogenous nucleotide sequences are included in a  
35 high copy number vector(s), it is preferred that the high copy number vector(s) is/are

selected from those that may be present at 20 to 500 (preferably, 400 to 500) copies per host cell. Particularly preferred high copy number vectors are the YEp vectors.

The method according to the invention enables the production of hydroxylated triple helical proteins. The term "triple helical protein" is to be understood as referring to a homo or heterotrimeric protein consisting of a polypeptide(s) or peptide(s) which include at least a region having the general peptide formula:  $(\text{GlyXY})_n$ , in which Gly is glycine, X and Y represent the same or different amino acids (the identities of which may vary from GlyXY triplet to GlyXY triplet) but wherein X and Y are frequently proline which in the case of Y becomes, after modification, hydroxyproline (Hyp), and n is in the range of 2 to 1500 (preferably 10 to 350), which region forms, together with the same or similar regions of two other polypeptides or peptides, a triple helical protein conformation. The triple helical proteins may include non-collagenous, non-triple helical domains at the amino and/or carboxy terminal ends or elsewhere.

Product-encoding nucleotide sequences used in the present invention encode a polypeptide or peptide of the general formula:  $(A)_l - (B)_m - [Z] - (C)_o - (D)_p$ , wherein; Z is a domain comprising two or more repeat units of the formula:  $[(E)_q - (\text{GlyXY})_i - (F)_r]$ , wherein E and F represent sequences of one or more amino acids (which sequences may vary from repeat unit to repeat unit), and for each repeat unit q and r are each independently selected from 0 and 1 and i is  $\geq 1$  such that domain Z comprises 2 to 1500 GlyXY triplets (preferably, 10 to 300 GlyXY triplets), Gly represents glycine, and X and Y (which may be the same or different), represent an amino acid, and wherein the identity of each amino acid represented by X and Y may vary from GlyXY triplet to GlyXY triplet, but wherein at least one Y of the  $(\text{GlyXY})_i$  sequence must be proline, A and D (which may be the same or different), each represent a polypeptide or peptide domain which optionally comprises a triple helical forming repeating sequence  $(\text{GlyXY})_n$ , and l and p are each independently selected from 0 and 1, B and C (which may be the same or different), each represent a polypeptide or peptide domain which is heterologous to collagen proteins and which does not comprise a triple helical forming repeating sequence  $(\text{GlyXY})_n$ , and m and o are each independently selected from 0 and 1.

Preferably, in domain Z, the component  $(\text{GlyXY})_i$  has an amino acid length which is at least three times greater than the combined amino acid length of the components E and F. Of course, in accordance with the formula of Z given above, one or both of E and F may be absent.

The portion of the product-encoding nucleotide sequence(s) encoding the repeat unit of domain Z may be generated through the use of polymerase chain reaction (PCR)

techniques or chemical DNA synthesis. Stepwise addition of such nucleotide sequences, so as to generate the repeating nucleotide sequence for domain Z may be achieved by utilising different restriction sites at the termini of primers used to produce the PCR fragments or through variations in chemical DNA synthesis. The selected restriction sequences are such that the desired linear order of the repeated nucleotide sequences is achieved in a manner which maintains the overall phase or open reading frame of the product-encoding nucleotide sequence and which ensures that every third amino acid of the encoded Z domain is Gly. Example 7 hereinafter provides an example of this strategy. That is, Example 7 describes a strategy for producing a product-encoding nucleotide sequence encoding a domain Z with three repeat sequences derived from an integrin binding site of Type III collagen, wherein the first step was to clone an EcoRI-[(GlyXY)<sub>n</sub>]-Bsp120I PCR fragment, the second step added a Bsp120I-[(GlyXY)<sub>n</sub>]-BssHII fragment, and the third step added a BssHII-[(GlyXY)<sub>n</sub>]-SacII, wherein the amino acid sequence of the polypeptide encoded by each fragment is the same. This strategy can be readily extended to add a nucleotide sequence encoding fourth, fifth etc. repeat units.

Alternatively, the portion of the product-encoding nucleotide sequence(s) encoding the repeat unit of domain Z may be generated using DNA ligase to join non-palindromic nucleotide sequences, which may be produced by PCR techniques or chemical DNA synthesis, end to end in such a manner as to maintain the open reading frame of the product-encoding nucleotide sequence and which ensures that every third amino acid of the encoded Z domain is Gly. The use of complimentary, but non-palindromic, overhanging sequences at the ends of the designed non-palindromic nucleotide sequences ensures that they are joined in a consistent head to tail orientation. Further, the strategy allows the ready linking of nucleotide sequences encoding other polypeptide or peptide domains (e.g. the abovementioned A, B, C and/or D), by utilising terminal non-palindromic overhang sequences which result in the generation of a restriction enzyme site. This restriction enzyme site can then be used for the additional cloning of nucleotide sequences encoding the other polypeptide or peptide domains.

This latter approach to the generation of the product-encoding nucleotide sequence(s) is preferred when the domain Z is to comprise a large number of repeat units (e.g.  $\geq 10$  repeat units).

The product-encoding nucleotide sequence(s) may include a sequence(s) encoding a secretion signal so that the polypeptide(s) or peptide(s) expressed from the product-encoding nucleotide sequence(s) are secreted.



The product-encoding nucleotide sequence(s) also comprise a nucleotide sequence preferably selected to match the codon use preferences of the selected yeast expression host, and is also constructed so as to minimise the potential for GGG to CCC interactions that may destabilize the structure.

5 Expression of the product-encoding nucleotide sequence(s) may be driven by constitutive yeast promoter sequences (e.g. ADH1 (Hitzeman *et al.*, 1981; Pihlajaniemi *et al.*, 1987), HIS3 (Mahadevan & Struhl, 1990), 786 (no author given, 1996 Innovations 5, 15) and PGK1 (Tuite *et al.*, 1982), but more preferably, by inducible yeast promoter sequences such as GAL1-10 (Goff *et al.* 1984), GAL7 (St. John & Davis, 1981), ADH2 (Thukral *et al.*, 1991) and CUP1 (Macreadie *et al.*, 1989).

10 The first and second nucleotide sequences encoding the P4H  $\alpha$  and  $\beta$  subunits can be of any animal origin although they are preferably of avian or mammalian, particularly human, origin (Helaakoski *et al.*, 1989). It is also envisaged that the first and second nucleotide sequences may originate from different species. In addition, the second nucleotide sequence encoding the P4H  $\beta$  subunit may include a sequence encoding an endoplasmic reticulum (ER) retention signal (e.g. HDEL (SEQ ID NO:13), KDEL (SEQ ID NO:42) or KEEL (SEQ ID NO:43)) with or without other target signals so as to allow expression of the P4H in the ER, cytoplasm or a target organelle or, alternatively, so as to be secreted.

20 Expression of the first and second nucleotide sequences may be driven by constitutive or inducible yeast promoter sequences such as those mentioned above. It is believed, however, that it is advantageous to achieve expression of the  $\alpha$  and  $\beta$  subunits in a co-ordinated manner using same or different promoter sequences with same induction characteristics, but preferably by the use of a bidirectional promoter sequence. Accordingly, it is preferred that the first and second nucleotide sequences be expressed by the yeast GAL1-10 bidirectional promoter sequence, although other bidirectional promoter sequences would also be suitable.

30 Multiple copies of the first, second and/or product-encoding nucleotide sequences may be introduced to the yeast host cell (e.g. present on a YAC vector or integrated into a host chromosome). It may be particularly advantageous to provide the product-encoding nucleotide sequence(s) in multicopy and, accordingly, it may be preferred to introduce the product-encoding nucleotide sequence(s) on a high copy number plasmid (e.g. a YEplasmid).

35 The introduced first, second and product-encoding nucleotide sequences may be borne on one or more stably retained and segregated DNA molecules. Where borne on more than one DNA molecule, the DNA molecules may be a combination of host

chromosome(s) and/or CEN sequence-including vector(s) in combination with high copy number vector(s). Some specific examples of yeast host cells suitable for use in the method according to the invention, are transformed with the following DNA molecules:

- 5 1. YEp-P3 + pYEUra3- $\alpha\beta$ ,
2. YEp-P3 + pYAC  $\alpha\beta$
3. YEpCEN-P3 + pYEUra3- $\alpha\beta$
4. YEpCEN-P3 + pYAC $\alpha\beta$
5. pYAC-P3 + pYAC  $\alpha\beta$
- 10 6. pYAC-P3 + pYEUra3- $\alpha\beta$
7. pYAC $\alpha\beta$ -P3;

wherein P3 represents a product-encoding nucleotide sequence(s),  $\alpha$  and  $\beta$  represent, respectively, nucleotide sequences encoding the P4H  $\alpha$  subunit and P4H  $\beta$  subunit, CEN represents an introduced centromere sequence. The pYEUra3 and pYAC vectors include CEN sequences.

Triple helical protein products produced in accordance with the method of the invention may be purified from the yeast host cell culture by techniques including standard chromatographic and precipitation techniques (Miller & Rhodes, 1982). For synthetic collagens, pepsin treatment and NaCl precipitation at acid and neutral pH may be used (Trelstad, 1982). Immunoaffinity chromatography can be used for constructs that contain appropriate recognition sequences, such as the Flag sequence which is recognised by an M1 or M2 monoclonal antibody, or a triple helical epitope, such as that recognised by the antibody 2G8/B1 (Glattauer *et al.*, 1997).

Yeast host cells suitable for use in the method according to the invention may be selected from genus including, but not limited to, *Saccharomyces*, *Kluveromyces*, *Schizosaccharomyces*, *Yarrowia* and *Pichia*. Particularly preferred yeast host cells may be selected from *S. cerevisiae*, *K. lactis*, *S. pombe*, *Y. lipolytica* and *P. pastoris*.

As indicated above, it is particularly preferred that the first, second and product-encoding nucleotide sequences be introduced to the yeast host cell by transformation with one or more YAC vectors. YAC vectors are linear DNA vectors which include yeast CEN sequences, at least one autonomous replication signal (e.g. *ars*) usually derived from yeast, and telomere ends (again, usually derived from yeast). They also generally include a yeast selectable marker such as URA3, TRP1, LEU2, or HIS3, and in some cases, an ochre suppressor (e.g. *sup4-o*) which allows for red/white selection in adenine requiring strains (i.e. the mutation of the adenine gene being due to a premature ochre stop codon). More commonly, two yeast selectable markers are

included, one on each arm of the artificial chromosome (each arm separated by the CEN). This allows selection of only those transformed hosts containing YACs with introduced sequences of interest within the desired restriction cloning site. That is, correct insertion of the sequences of interest (e.g. an expression cassette) rejoins the two arms of the restricted YAC, thus rendering transformants prototrophic for both markers. YACs have been designed to allow for the introduction of large exogenous nucleotide sequences (i.e. of the order of 100kb or more) into yeast host cells. The present inventors have hereinafter shown that such YACs may be used for the stable expression of multiple exogenous nucleotide sequences (e.g. nucleotide sequences encoding a natural collagen and both the  $\alpha$  and  $\beta$  subunits of P4H).

In some embodiments of the invention, it may be preferred that one or more (but not all) of the first, second and product-encoding nucleotide sequences be introduced to the yeast host cell by transformation with one or two YE<sub>p</sub> vectors. YE<sub>p</sub> vectors carry all or part of the yeast 2 $\mu$  plasmid with at least the *ori* of replication. They also include a yeast selectable marker such as HIS3, LEU2, TRP1, URA3, CUP1 or G418 resistance, and often also contain a separate *ori*, generally ColE1, and markers, such as ampicillin resistance, for manipulation in *E.coli*. They show high copy number, for example 20-400 per cell, and are generally efficiently segregated. Stability during cell division is dependent on the vector also containing the REP2/STB locus from the 2 $\mu$  plasmid. However, stability is not as good as endogenous 2 $\mu$  plasmid of the host, particularly when heterologous genes are induced for expression. Stability also declines with increasing plasmid size. (Wiseman, 1991).

The method of the present invention enables the production of triple helical protein products with two or more repeat units, which allows control of biological function and permits the possibility of enhancing the efficacy of selected domains by, for example, increasing binding sites and avidity for interacting agents and by activation through receptor clustering.

The terms "comprise", "comprises" and "comprising" as used throughout the specification are intended to refer to the inclusion of a stated component or feature or group of components or features with or without the inclusion of a further component or feature or group of components or features.

The invention will now be described by way of reference to the following non-limiting examples and accompanying figures.

**Brief description of the accompanying figures:**

Figure 1 shows, diagrammatically, the construction of the expression vector pYEUra3.2.129#39I#5 (labeled pYEUra3-M $\beta\alpha$ ).

5 Figure 2 shows the nucleotide sequence for the COLIII1.6 kb DNA (SEQ ID NO:39).

Figure 3 shows, diagrammatically, regions of the human collagen III gene that have been isolated by PCR. The 1.6kb DNA used in the examples hereinafter is also shown. It is to be understood that the other regions shown in the figure could substitute for the COLIII1.6kb DNA in those examples.

10 Figure 4 shows, diagrammatically, the construction of the expression vector YEpFlagCOLIII1.6kb (labeled YEpFlag-C3).

Figure 5 shows, diagrammatically, the construction of pYAC5  $\beta\alpha$ .

Figure 6 shows, diagrammatically, the construction of pYAC  $\beta\alpha$ -COL III1.6 kb.

Figure 7 outlines the construction of synthetic collagen products.

15 Figure 8 provides the nucleotide sequence (SEQ ID NO:40) for SYN-C3 together with the amino acid sequence (SEQ ID NO:41) of the encoded polypeptide.

**Example 1: Construction of a yeast vector for co-ordinated co-expression of the  $\alpha$  and  $\beta$  subunits of Prolyl-4-hydroxylase.**

20 *Production of yeast expression vector:*

pYEUra3 (Clontech) contains the bidirectional promoter for GAL1-10 expression. Induction by galactose in the absence of glucose results in high level expression from pGAL1 of any protein encoded by DNA sequences inserted in the correct orientation in the MCS (multiple cloning site) [either XhoI, Sall, XbaI or  
25 BamHI sites] provided there is an initiating ATG start codon. For pGAL10, expression induced by galactose occurs if the DNA sequences to be expressed are inserted in frame with the ATG codon of GAL10 when said DNA sequences to be expressed is inserted in the EcoRI site.

In order to utilise the EcoRI site for cloning, without the necessity that the insert  
30 be in frame with the ATG of GAL10 for expression, it was necessary to modify pYEUra3 to remove the GAL10 initiation codon. This was done as follows. A PCR fragment was generated using pYEUra3 as template and primers 3465 [5'CTG.TAG.AGG.ATC.CCCGGG.TAC.GGA.GC-3' (SEQ ID NO:1), where the BamHI site is underlined] and primer 1440  
35 [5'TTA.TAT.TGA.ATT.CTC.AAA.AAT.TC-3' (SEQ ID NO:2), where the EcoRI restriction site is underlined]. Primer 1440 introduces an EcoRI site preceding the

initiating ATG of GAL10 in pYEura3. The PCR fragment was restricted with BamHI and EcoRI and cloned into pYEura3 similarly digested with BamHI and EcoRI, replacing the BamHI-EcoRI fragment containing an ATG start codon with a BamHI-EcoRI fragment lacking this ATG, to generate plasmid pYEura3.2.12. The EcoRI site  
 5 can then be used as a cloning site for which an initiating codon must be provided by the inserted DNA sequence as with the MCS at the other end of the promoter, thus placing it under control of the bidirectional pGAL1-10 promoter and rendering expression inducible by galactose as are DNA sequences inserted in the MCS at the other end of the promoter. Cloning DNA sequences in the MCS and in the EcoRI site allows for co-  
 10 ordinate expression by the bidirectional promoter when induced by galactose.

*Isolation of DNA molecules encoding the  $\alpha$  and  $\beta$  subunits of P4H:*

The  $\alpha$  subunit of P4H was PCR amplified from cDNA (Clontech Human Kidney Quick Clone™ cDNA Cat.#7112-1) using primers 1826 [5'-TGT.AAA.ATT.AAA.GGA.TCC.CAA.AG.ATG.TGG.TAT-3'] (SEQ ID NO:3), where the BamHI  
 15 site is underlined, ATG is the initiating codon for  $\alpha$  subunit] and 1452 [5'-GCCG.GGA.TCC.TG.TCA.TTC.CAA.TGA.CAA.CGT-3'] (SEQ ID NO:4), wherein the BamHI site is underlined, TCA is the translation stop codon]. Two isoforms were obtained and cloned into the BamHI site of pBluescript II SK+ [Stratagene Cat.#212205] as storage vector to give pSK+ $\alpha$ .1 (form I) and pSK+ $\alpha$ .2 (form II). There are  
 20 no BamHI sites in the DNA encoding the  $\alpha$  subunit. The signal sequence for secretion is present in the BamHI fragment of both forms.

The  $\beta$  subunit of P4H [also known as PDI/protein disulfide isomerase] [Pihlajaniemi *et al.*, 1987] was PCR amplified from cDNA (Clontech Human Kidney Quick Clone™ cDNA Cat.#7112-1) using primer pairs 2280 [5'-  
 25 AC.TGG.ACG.GAT.CCC.GAG.CGC.CCC.GCC.TGC.TCC.GTG.TCC.GAC.ATG-3'] (SEQ ID NO:5)] and 2261 [5'-G.GTT.CTC.CTT.GGT.GAC.CTC.CCC.TT-3'] (SEQ ID NO:6), where the BstEII site is underlined] for the amino terminal part of the  $\beta$  subunit and primer pairs 2260 [5'-GAA.GGG.GAG.GTC.ACC.AAG.GAG.AAC-3'] (SEQ ID NO:7), where the BstEII  
 30 site is underlined] and 1932 [5'-CC.TTC.AGG.ATC.CTA.TTA.GAC.TTC.ATC.TTT.CAAC.AGC-3'] (SEQ ID NO:8)] for the carboxy terminal part of the  $\beta$  subunit. The two PCR fragments for the  $\beta$  subunit were then ligated together following BstEII digestion, to produce a single fragment encoding the entire  $\beta$  subunit. This fragment was then amplified using the  
 35 primers 2280 [5'-AC.TGG.ACG.GAT.CCC.GAG.CGC.CCC.GCC.TGC.TCC.

GTC.TCC.GAC.ATG-3' (SEQ ID NO:9), where the BamHI site is underlined, and ATG is the initiating codon for the  $\beta$ -subunit] and primer 1932 [5'-CC.TTC.AGG.ATC.CTA.TTA.GAC.TTC.ATC.TTT.CAC.AGC-3' (SEQ ID NO:10), where the BamHI site is underlined, and TTA is the translation stop codon for the  $\beta$  subunit] and then cloned

5 into the BamHI site of pBluescript SKII+ to generate the storage vector pSK+ $\beta$ . Subsequently, the BamHI fragment of pSK+ $\beta$  was amplified by using primers 2698 [5'-CTA.GTT.GAA.TTC.TAC.ACA.ATG.CTG.CGC.CGC.GCT. CTG.CTG-3' (SEQ ID NO:11), where the EcoRI site is underlined, and ATG is the initiating codon of the  $\beta$  subunit] and 2699 [5'-GCA.ATG.GAA.TTC.TTA.TTA.

10 CAG.TTC.GTG.CAC.AGC.TTT-3' (SEQ ID NO:12), where the EcoRI site is underlined, and TTA.TTA. provides two translation stop codons, and GTG. changes a lysine [K] residue to a histidine [H] residue to provide a native yeast ER retention signal, HDEL (i.e. His.Asp.Glu.Leu (SEQ ID NO:13)) rather than a mammalian KDAEL (SEQ ID NO:14) ER retention signal]. The resultant PCR fragment was then

15 blunt end cloned into the SrfI site of pCRScript [Stratagene, Cat.# 211190] to generate pCRScript $\beta$ . After retrieving the EcoRI fragment containing the  $\beta$  subunit from pCRScript $\beta$  by EcoRI digestion, the fragment was again cloned into the EcoRI site of pCRScript to generate pCRScript $\beta$ EcoRI#4.

*Construction of yeast expression vector including fragment encoding the  $\alpha$  and  $\beta$  subunit of P4H:*

20 The  $\beta$  subunit fragment was obtained as an EcoRI fragment from EcoRI digestion of pCRScript $\beta$ EcoRI#4. This EcoRI fragment was cloned into the EcoRI site of pYEUra3.2.12 to generate plasmid pYEUra3.2.12 $\beta$ #39. The  $\alpha$  subunit fragment from pSK+ $\alpha$ .1 was re-excised from pSK $\alpha$ .1 by BamHI and cloned into the BamHI site

25 of pYEUra3.2.12 $\beta$ #39 to give pYEUra3.2.12 $\beta$ #39 $\alpha$ #5] (Figure 1). The  $\beta$  subunit fragment is under control of pGAL10 and the  $\alpha$  subunit fragment is under control of pGAL1. This is a bidirectional promoter and allows co-ordinated induced expression of both subunits of prolyl-4-hydroxylase. Both fragments provide a native ATG initiating codon for translation. The encoded  $\beta$  subunit has its own signal secretion

30 signal and a HDEL endoplasmic retention (ER) sequence at the carboxy terminus of the protein. While the encoded  $\alpha$  subunit with its own signal sequence has no ER retention signal it should, nevertheless, be retained through its interaction with the  $\beta$  subunit.

**Example 2: Co-ordinated co-expression of a collagen segment and prolyl-4-hydroxylase ( $\alpha$  and  $\beta$  subunit) and synthesis of hydroxylated collagen Type III in yeast.**

A 1.6 kbp recombinant collagen fragment was generated by PCR using primers 1989 [Forward primer 5'-gct.agc.aag.ctt GGA.GCT.CCA. GGC.CCA.CTT.GGG.ATT.GCT.GGG-3' (SEQ ID NO:15)] and 1903 [Reverse primer 5'-tcg.cga.tct.aga.TTA.TAA.AAA.GCA.AAC.AGG.GCC.AAC.GTC.CAC. ACC-3' (SEQ ID NO:16)] homologous to a region of the collagen type III alpha I chain (COL3A1). The template for isolation of the fragment of type III collagen alpha 1 chain was prepared from Wizard purified DNA obtained from a cDNA library [HL1123n Lambda Max 1 Clontech Lot#1245, Human Kidney cDNA 5'-Stretch Library].

The actual size of the isolated 1.6 kbp fragment is 1635 bp, comprising 1611 bp of COL3A1 DNA flanked either side by 12bp derived from the primers. The 1611 bp of COL3A1 DNA corresponds to nucleotides #2713-4826 (i.e codon #905-1442) of the full-length coding sequence, thereby spanning a portion of the I-helix region, all of the C-telo-peptide, all of the C-pro-peptide and stop codon.\*<sup>1</sup> The nucleotide sequence for the COL3A1 DNA is provided at Figure 2. The region covered by the COL3A1 DNA is shown at Figure 3. The 1.6kbp fragment has a NheI [GCTAGC (SEQ ID NO:17)] site and a HindIII [AAGCTT (SEQ ID NO:18)] site added at the 5'-end and a XbaI [TCTAGA (SEQ ID NO:19)] site and a NruI [TCGCGA (SEQ ID NO:20)] site added at the 3' end [where the 5' end is taken to be the forward direction of the reading frame, ie the amino terminal end of the derived coding sequence, and the 3' end is that derived from the reverse primer corresponding to the 3' end of the gene and carboxy end of the derived amino acid sequence]. This confers portability on the collagen fragment.

The 1.6kbp fragment was cloned into the SmaI site of YEplFlag1 [IBI Catalogue #13400] so that the coding sequence is fused in frame with the vector expressed Flag protein. This allows for in frame expression of the introduced collagen gene fragment as a fusion protein when grown on ethanol. The blunt end cloning was performed by ligation of the SmaI digested vector sequence [gel purified] and the 1.6kbp PCR fragment [gel purified, non-phosphorylated] at 20°C, in the presence of SmaI, to prevent recircularisation of the vector alone and reduce the level of false positive transformants obtained. There are no SmaI, NheI, HindIII, XbaI or NruI sites in the fragment of collagen DNA used in the cloning.

Small scale mini-preparations [prepared using Bio101 columns and described methods for their use] of DNA from ampicillin resistant transformant colonies of *E.coli* were screened by restriction enzyme analysis. 10ml cultures rather than 1 ml cultures

were required to prepare an adequate level of DNA for analysis, as YEpFlag plasmids do not appear to be at a high copy number in *E.coli*.

The fusion protein was of the form: yeast  $\alpha$  factor signal sequence for direction to the ER and commitment to the yeast secretion pathway, yeast  $\alpha$  factor propeptide with cleavage sites for kex 2-endopeptidase, resulting in removal of all  $\alpha$ -factor amino acid residues and generation of a free Flag-tagged amino terminal end, Flag peptide for detection and tagging of the fusion protein (8 amino acid residues), linker peptide (4 amino acid residues), collagen helix (255 amino acid residues), collagen C-telopeptide [C-tel] (25 amino acid residues) and C-propeptide [C-pro] (255 amino acid residues) (for aid in formation of triple helix). The expected Flag-tagged protein consists of 547 amino acid residues with a expected MW of ~60kDa].

Expression of the fusion protein in YEpFlag1 is under the control of the ADH2 promoter which is repressed by glucose but active in the presence of ethanol [a by-product of glucose metabolism]. There are multiple copies of the vector in individual yeast transformants due to the presence of the yeast 2 micron origin of replication in the vector, which leads to elevated expression of the 1.6 kbp PCR collagen fragment when glucose repression is lifted by consumption of glucose during growth. One unique feature of this cloning scheme is that inserts of the 1.6kbp collagen fragment in the wrong orientation will not form fusion products as the terminal leucine residue preceding the stop codon is coded by the codon AAT. In reverse orientation this generates a stop codon TAA. The result of incorrect insertion is the addition of only a single leucine coding codon [the stop codon TAA in reverse is AAT] following the Flag sequence before the protein is terminated.

The amino acid sequence of the Flag-tagged fusion protein at the point of fusion is N-Asp-Tyr-Lys-Asp-Asp-Asp-Asp-Lys-[Flag]-Ala-Ser-Lys-Leu-[linker]-Gly-Ala-Pro-Gly-Pro-Leu-Gly-Ile-Ala-[ $\alpha$ -helix] (SEQ ID NO:21).

The YEpFlag collagen construct [hereinafter referred to as YEpFlag COLIII1.6kb; Figure 4] was introduced into a tryptophan prototrophic yeast strain such as for example BJ3505 [a pep4::HIS3 prb-1.6R HIS3 lys2-208 trp1--101 ura3-52 gal2 can1], BJ5462 [ $\alpha$  ura3-52 trp1 leu2-1 his3-200 pep4::HIS3 prb-1.6R can1 GAL], (YGSG) JHRY1-5D $\alpha$  [ $\alpha$  his4-519 ura3-52 leu2-3 leu2-112 trp1 pep4-3] or KRYD1 [BJ3505xBJ5462 diploid] by transformation using electroporation, lithium acetate or spheroplast regeneration. Tryptophan auxotroph transformants were obtained, grown to high cell density in selective media [lacking tryptophan] followed by transfer to YPHSM, YEPM or YEPD or YEPGal, YEPE as described in the protocol provided with the YEpFlag expression system [IBI catalogue #13400]. At 3-9 days following



inoculation 1ml aliquot's of culture were made and pellets and supernatants separated by centrifugation at 13000rpm in a benchtop centrifuge. Total yeast pellets were resuspended in 100µl of gel loading buffer [5xSDS] containing PMSF [0.002M], vortexed vigorously for 2 minutes, and boiled for 5 minutes. From the pellets 900µl supernatants were retained to which 100µl 5xSDS/0.002M PMSF was added, and treated as described for the pellets. For both pellets and supernatants 20µl aliquot's were assayed by Western blot analysis of SDS-PAGE yeast total protein or of supernatants [media] following transfer to nitrocellulose and prehybridisation of the filters in blotto. Western blotting was carried out using α-Flag MAb M1 [against N-terminal free Flag] (International Biotechnologies Inc., (Eastman Kodak) Cat. No. IB13001) or M2 [against Flag] (International Biotechnologies Inc., (Eastman Kodak) Cat. No. IB13010).

Western blots revealed the presence of a protein band of approximately 60kDa. This is the expected size of a protein fusion containing Flag-helix-C-tel-C-pro. After prolonged incubation the Flag responsive antibodies detected the appearance of the fusion product in the media. Detection in both pellet and media supernatant with M1 antibody demonstrates that the α factor leader has been completely removed. No precursor forms with α factor pro-region [glycosylated or not] were observed.

No band corresponding to 60kDa was obtained which hybridised to M1 or M2 with proteins obtained from untransformed yeast hosts. When yeast transformed with YEpFlag [no insert] alone was used, bands were obtained in pellets, but only with M2 MAb. These bands correspond to un-secreted α-proregion-with C-terminal Flag and various glycosylated forms of the same. No Flag is detected in supernatants but this is to be expected as it is only 8 amino acids long. No expression from the ADH2 promoter for any construct is observed in the presence of glucose.

YEpFlagCOLIII 1.6kb was also co-introduced [co-transformed] into yeast strains such as BJ5462 and KR DY1 which are capable of growth on galactose along with pYEUra3 [Clontech] [pYEUra3 and its derivatives contain the bidirectional GAL1-10 promoter. Both the ADH2 and GAL1-10 promoters are repressed by glucose. The GAL1-10 promoter is induced by galactose] or pYEUra3.2.12 [a modification of the Clontech parent vector which allows cloning of genes into an EcoRI site without the necessity of the introduced gene being in the correct reading frame] or pYEUra3.2.12β#39 [in which the DNA encoding the β subunit (equivalent to protein disulfide isomerase of prolyl-4-hydroxylase is cloned into the EcoRI site of pYEUra3.2.12 under control of GAL10 promoter] or pYEUra3.2.12β#39α#5 [in which

the DNA encoding the  $\alpha$  subunit of P4H is cloned into the BamHI site of pYEUra3.2.12 $\beta$ #39 under control of the GAL1 promoter].

Transformants were selected on media lacking tryptophan or uracil or lacking both tryptophan and uracil. As previously done with tryptophan transformants obtained above with YEpFlag or YEpFlagCOLIII1.6kb, transformants were grown in selective media prior to growth in YPHSM, YEPM, YEPD, YEPG or YEPE and after 4 days galactose was added to a final concentration of 2%, 0.5% or 0.2%. Total yeast protein or supernatants were analysed by Western blot analysis as described above except that a third MAb [5B5 against the  $\alpha$  subunit] (Dako Corporation, Cat. No. M877) was also used.

Western blot analysis revealed the presence of a ~60kDa band in  $trp^-$  or  $trp^- ura^-$  yeast transformed with YEpFlag COLIII1.6kb but not YEpFlag alone when screened with MAb M1 or M2 as was previously the case with transformants obtained with single plasmid transformation.

Analysis also showed the presence of a ~60kDa band in  $ura^-$  or  $ura^- trp^-$  but not  $trp^-$  yeast transformants transformed with pYEUra3.2.12 $\beta$ #39 or pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 or cotransformed with same plus YEpFlag or YEpFlagCOLIII 1.6kb when screened with anti- $\beta$  subunit MAb 5B5 but only following induction with galactose and only when galactose was between 0.2 and 0.5% and not at 2%. The expected size for the  $\beta$  subunit is also 60kDa. This band is not detected by M1 or M2 in uracil auxotrophic yeast transformed with pYEUra3.2.12 $\beta$ #39 or pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 alone.

At the time of the experimentation, an antibody for the detection of expression of the  $\alpha$  subunit from the bidirectional GAL1,10 promoter in pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 was not available but as the promoters for both GAL1 and GAL10 are normally co-induced and under the control of the same UAS (upstream activation sequence) in yeast it was assumed that the  $\alpha$  subunit is also transcribed and expressed where the  $\beta$  subunit is demonstrated to be expressed. To test this, the capacity for pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 / YEpFlag COLIII 1.6kb co-transformants induced with 0.2% galactose following at least 4 days growth on YPHSM to produce functional P4H was examined. Galactose was added following the clear demonstration of the expression of Flag-collagen by a positive response of yeast protein to M1 or M2 in Western blots and the absence of a response to MAb 5B5 against  $\beta$  subunit. Following induction with galactose [16hrs] protein was again examined and the presence of M1 or M2 responsive bands and 5B5 responsive bands were separately demonstrated. Protein was transferred to PVDF membrane following SDS-PAGE and the membrane sliced into strips. Membrane

strips containing protein from the region corresponding to the 60kDa responsive area was subject to hydrolysis and amino acid analysis. Amino acid analysis revealed the presence of hydroxyproline in this material from co-transformants of yeast co-transformed with YEpFlagCOLIII1.6kb and pYEUra3.2.12β#39α#5 after induction with 0.2% galactose but no hydroxyproline was detected with protein from control samples with or without galactose.

The media used contains peptone derived from bovine protein hydrolysates but no hydroxyproline was found in total yeast grown on this media nor in any of the singly transformed yeast [one vector alone]. Only in yeast co-transformants was hydroxyproline detected in the 60kDa bands and then only when galactose was added. Uninduced co-transformants [no galactose] in which Flag detected collagen was expressed did not contain any hydroxyproline in the 60kDa band excised from PVDF following transfer. Hydroxyproline was only found in the 60kDa region and not in other regions of the blot.

The clear evidence then, is that following galactose induction of pYEUra3.2.12β#39α#5 a product is produced in yeast which is capable of hydroxylating the proline residues of a co-expressed Flag-tagged collagen fragment. Such activity is not found in yeast untransformed or transformed with pYEUra3.2.12β#39 [no α subunit] or in uninduced yeast grown on ethanol or glucose.

A clear advantage of this method of co-expression for the production of hydroxylated collagen in yeast is the co-ordinated expression of the three genes that is possible in co-transformants. Another advantage is that the α and β subunits themselves are co-ordinately expressed. A third advantage is that the αβ expression vector (i.e. pYEUra3.2.12β#39α#5) contains a centromere sequence and behaves as a mini-chromosome. It is therefore very stable and does not require selection pressure to be maintained for its stability. The removal of selection pressure in yeast does not appear to effect the stability of the YEpFlag collagen construct as it is in very high copy number, but clearly the ability to only be concerned with maintenance of a single plasmid in the absence of selection pressure is important rather than balancing the effects of selection pressure on the stability of three separate plasmids if the α, β and collagen fragments were separately cloned on multicopy vectors. Also the use of a bidirectional promoter to express the α and β subunits simultaneously is of benefit rather than expressing them from different promoters on different plasmids in different amounts. The α subunit probably requires the synthesis of equal or higher levels of the β subunit for its correct assembly into functional P4H (α<sub>2</sub>β<sub>2</sub>) enzyme and co-ordinated expression appears to be an efficient mechanism to ensure this.

\*<sup>1</sup> [ Codon numbering for collagen type III alpha 1 chain: ATG, codon #1; codon #1-codon #24, signal sequence; codon #25-codon #116, N-pro-peptide sequence, codon #117-codon #130, N-telo-peptide sequence; codon #131-codon #1161,  $\alpha$ -helix sequence; codon #1162-codon #1186, C-telo-peptide; codon #1187-codon #1441, C-pro-peptide; codon #1442, stop] and [corresponding nucleotide numbering for collagen type III alpha 1 chain: nucleotide #1-72, signal sequence; nucleotide #73-348, N-pro-peptide sequence; nucleotide #349-390, N-telo-peptide; nucleotide #391-nt#3983,  $\alpha$ -helix region; nucleotide #3984-4058, C-telo-peptide; nucleotide #4059-4823, C-pro-peptide sequence; nucleotide #4824-4826, stop codon].

### **Example 3: Use of Yeast Artificial Chromosomes [YACs] for co-ordinated expression of the $\alpha$ and $\beta$ subunits of Prolyl-4-hydroxylase [P4H].**

pYAC5 [11454bp] (Kuhn and Ludwig, 1994) was digested with BamHI to liberate the HIS3 gene [1210bp] from between the 2 telomere ends and with Sall-NruI to produce two fragments [left arm: fragment 1, 5448bp & right arm: fragment2, 4238bp] which were gel purified. Fragment 1 was BamHI-telomere end-*E.coli* ori- $\beta$ -lactamase gene [ampicillin-resistance] -TRP1-ARS1-CEN4-tRNAsup-o-Sall. Fragment 2 was BamHI-telomere end-URA3-NruI.

pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 was digested with Sall-EcoRV to produce a P4H expression cassette fragment of the form Sall-XbaI-BamHI- $\alpha$ -ATG-BamHI-pGAL1-10-EcoRI-ATG- $\beta$ -EcoRI-SmaI-EcoRV [4864bp] which was gel purified. The expression cassette fragment encoding the  $\alpha$  and  $\beta$  subunits of P4H under the control of a galactose inducible bidirectional promoter was ligated with fragments 1 and 2 of the BamHI-Sall-NruI digested pYAC5 and the ligation mix used to transform the following yeast strains: BJ2407 [a/ $\alpha$  prb1-11222/prb1-1122 prc1-407/prc1-407 pep4-3/pep4-3 leu2/leu2 trp1/trp1 ura3-52/ura3-52 ], KRYD1 [a/ $\alpha$  ura3-52/ura3-52 trp1- $\Delta$ 101/trp1 lys2-208/LYS2 HIS3/his3 $\Delta$ 200 gal2/GAL2 can1/can1 pep4::HIS3/pep4::HIS3 prb1 $\Delta$ 1.6R/prb1.6R ], GY1 [  $\alpha$  leu2 ade1 trp1 ura3 ], JHRY1-5D $\alpha$  [  $\alpha$  his4-519 ura3-52 leu2-3 leu2-112 trp1 pep4-3 ], and YPH150[  $\alpha$ /a ura3-52/ura3-52 lys2-801a/lys2-801a ade1-101o/ade1-101o leu2 $\Delta$ 1/leu2 $\Delta$ 1 trp1- $\Delta$ 63/trp1- $\Delta$ 63 his3 $\Delta$ 200/his3 $\Delta$ 200 ] using the method for lithium acetate transformation. Yeast strains were also transformed with pYAC5 digested with BamHI and undigested pYAC5.

Ura<sup>+</sup> Trp<sup>+</sup> co-transformants were obtained for all strains where the two fragments of pYAC5 each carrying either TRP1 [Sall-CEN4-TRP1-BamHI] [fragment 1] or URA3 [NruI-URA3-BamHI] [fragment 2] as the selectable marker for transformation each on one arm of the YAC, had been linked together by the insertion

of the P4H expression cassette into the Sall-EcoRV sites. This vector was designated pYAC5 $\beta\alpha$  (Figure 5). The vector was of the form BamHI-telomere-URA3-NruI/EcoRV [both sites destroyed]- $\beta$ -ATG-pGAL10-1-ATG- $\alpha$ -Sall-tRNA<sup>sup</sup>-CEN4-ARS1-TRP1-AMPr-ori-telomere-BamHI. The presence of the CEN4 sequence means  
 5 the vector behaves as a stable chromosome during replication and is segregated at least 1 copy per cell at mitosis and meiosis [as was the case for pYEura3.2.12 $\beta$ #39 $\alpha$ #5]. The telomere ends mean that the vector is linear and stable.

Transformants and controls [pYAC5 alone (circular), pYAC5 linearised by BamHI digestion] were replica plated onto nitrocellulose filters laid over selective  
 10 media [SD Complete lacking uracil and tryptophan] or rich media [YEpd] and incubated 2-5 days at 30C till confluent. Filters were transferred to selective media containing galactose [2%] instead of glucose or rich media containing galactose [2%] as well as glucose media plates and grown at 30C for periods between 2h-72h. At the end of incubation colonies were lysed on 0.1%SDS-0.2N NaOH-0.1%  $\beta$ -  
 15 mercaptoethanol, washed with water and filters blocked with Blotto. Production of the  $\alpha$  and  $\beta$  subunits of P4H was ascertained by hybridising the treated filters with MABs specific for the  $\alpha$  [MAB 9-47H10] (ICN Biomedical Inc. Cat. No. 631633) and  $\beta$  [MAB 5B5] subunits. Colonies transformed with pYAC5 $\beta\alpha$  and induced with galactose showed hybridisation with MABs against the subunits of P4H demonstrating co-  
 20 ordinated production of  $\alpha$  and  $\beta$  from the bi-directional GAL 1-10 promoter. Controls filters and control yeast did not produce a response to P4H MABs. Yeast transformants carrying pYAC5 $\beta\alpha$  grown on glucose [a repressor of the bi-directional GAL 1-10 promoter] also did not produce a positive response.

Positive transformants identified in the above screening procedure were  
 25 precultured/grown in 10ml liquid culture media containing selective media lacking ura and trp or rich media [containing glucose, glycerol or raffinose]. Aliquots were transferred to inducing media [selective or rich] containing 0.2-2% galactose. Where glucose was the carbon source pellets were washed in sterile water prior to induction. After 2-20h further growth at 30C cell pellets were collected, suspended in loading  
 30 buffer and total yeast protein separated on SDS-PAGE and western blotted. Filters were blocked with blotto and hybridised with MABs against both of the P4H subunits. Only those yeast transformants carrying pYAC5 $\beta\alpha$  and induced with galactose gave the expected 60kDa bands for  $\alpha$  and  $\beta$  subunits. This demonstrates that the P4H expression cassette has been functionally inserted into pYAC5. The advantage of  
 35 having the P4H cassette in the pYAC is twofold; [1] as with the case of pYEura3.2.12 $\beta$ #39 $\alpha$ #5 the presence of the CEN sequence means that the vector is

stably maintained in this system when selection pressure is removed for growth in rich media, which increases yield through increased cell density, and [2] the pYAC5 $\beta\alpha$  construct allows for the subsequent insertion of multiple and different triple helical protein expression cassettes.

5

**Example 4: Co-expression of collagen/triple helical protein fragment(s) expressed on a multicopy plasmid and P4H subunits in yeast transformants carrying pYAC5 $\beta\alpha$ .**

Yeast host strains containing pYAC5 $\beta\alpha$  or pYAC5 were transformed with  
 10 YEpFlagColIII 1.6kb or YEpFlag alone. The form of the collagen bearing vector was circular and multicopy. In this instance, as the YEpFlagCOLIII 1.6kb and the pYAC constructs both contain the same selectable marker, yeast transformants producing Flag tagged-collagen were identified by colony hybridisation with MAb's against Flag [M1 or M2]. Colonies were also screened for whether they carried extra copies of *bla* gene  
 15 [multicopy] by identifying those colonies producing increased levels of  $\beta$ -lactamase by PADAC assay (Macreadie *et al.*, 1994). In other examples, the multicopy plasmid could utilise a different selectable marker other than URA3 or TRP1 found on each arm of the YAC. Various co-transformant types carrying pYAC5 $\beta\alpha$  and YEpFlag COLIII 1.6kb were assayed as in Example 1 for collagen production, P4H subunit production,  
 20 and P4H activity. Those co-transformants containing pYAC5 $\beta\alpha$  plus YEpFlag COLIII 1.6kb were then screened as described in the previous example for hydroxylated collagen to identify 60kDa bands in western blots responding to MAb's against the  $\alpha$  and  $\beta$  and Flag following induction. The  $\alpha$  and  $\beta$  subunits were only identified following galactose induction. Hydroxylated protein was only identified following  
 25 induction of both the  $\alpha$  and  $\beta$  subunits of P4H.

**Example 5: Introduction of collagen expression cassette into pYAC5 and pYAC5 $\beta\alpha$ .**

YEpFlag was linearised by digestion with ScaI which cuts at a single  
 30 recognition site in the ampicillin resistance gene for  $\beta$ -lactamase [*bla*]. There are no ScaI sites in the 1.6kb collagen fragment insert so ScaI could also be used to linearise YEpFlagColIII 1.6kb. Linear DNA was used to transform yeast containing pYAC5 or pYAC5 $\beta\alpha$ . Yeast transformants producing Flag tagged-collagen were identified by colony hybridisation with MAb's against Flag [M1 or M2]. Colonies carrying extra  
 35 copies of *bla* gene [multicopy] were also identified. Those colonies producing increased levels of  $\beta$ -lactamase by the PEDAC assay were found to have inserted a

copy of YEpFlag COLIII 1.6kb into the pYAC5 or pYAC5 $\beta\alpha$  vector of the host strain and correspond to those colonies positive to MAbs M1 or M2. The increased  $\beta$ -lactamase activity is a result of gene amplification resulting from homologous recombination between the linearised *bla* gene on YEpFlagCOLIII 1.6kb and the *bla* gene on pYAC. The new plasmids formed by insertion into pYAC5 or pYAC5 $\beta\alpha$  of the YEpFlag COLIII 1.6kb vector were designated pYAC-COLIII 1.6kb and pYAC  $\alpha\beta$ -COLIII 1.6kb (Figure 6). Expression experiments were performed and only those strains carrying all 3 genes on the YAC [pYAC  $\beta\alpha$  -COLIII 1.6kb] and induced for P4H with galactose produced hydroxylated collagen.

#### **Example 6: Cloning and expression of a synthetic collagen protein.**

A strategy is described for the generation of "synthetic/novel" collagen proteins involving the *in vitro* assembly of synthetic oligonucleotides repeat sequences encoding the peptides GPP.GPP.GLA (SEQ ID NO:22), GPP.GPP.GER (SEQ ID NO:23), GPP.GPP.GPA (SEQ ID NO:24) or GPP.GPP.GAP (SEQ ID NO:25). The synthetic collagen sequences are engineered to contain a high percentage of proline residues as this residue has been shown to confer thermal stability to collagen molecules. The residue pairs chosen in the above peptides for the XY position (i.e. LA, ER, PA or AP), are selected since they appear in statistically higher amounts in fibrillar collagens.

Mixtures of synthetic oligonucleotides encoding SEQ ID NOs:22, 23, 24 or 25 may be joined together to generate DNA fragments of discrete lengths, encoding synthetic collagen proteins of discrete molecular size and with different physical characteristics. These synthetic gene segments can be cloned into various expression vectors for subsequent production of a collagen product in yeast. An outline of the strategy for construction of a synthetic oligonucleotide encoding a collagen is shown in Figure 7 where XY is shown, for the purposes of exemplification only, as ER, LA, AP, PA.

Such synthetic oligonucleotides have been synthesised and several libraries containing gene segments of various lengths have been generated by ligating these oligonucleotides together (maximum visible DNA length approx. 1000 base pairs coding for a polypeptide of ~ 350 amino acid residues).

#### **Example 7: Construction of a synthetic hydroxylated triple helical protein for stable expression in yeast.**

A region of Type III collagen was selected for its known capacity to bind and activate platelets [through an integrin binding site near -Gly-Leu-Ala-Gly-Ala-Pro-Gly-

Leu-Arg (SEQ ID NO:26)]. A region of 5 GLY-X-Y repeats to the N-terminal side and 7 GLY-X-Y repeats to the C-terminal side were also included to form the basic repeat unit for inclusion in the synthetic fragment. The sequence of the repeat was GGKGDAGAPGERGPP-GLAGAPGLR-GGAGPPGPEGGKGAAGPPGPP (SEQ ID NO:27). This corresponds to residues 637-681 (nucleotides 1909-2043) in the COL3A1 gene [with Met =1]. At the 5'-end of the DNA an EcoRI site and NheI site was included such that the NheI site provided an initiating methionine. Thus the sequence at the amino end is MGAPGAP (SEQ ID NO:28), where GAPGAP (SEQ ID NO:29) is the natural sequence flanking the repeat in COL3A1. The repeat was linked to a second repeat by a linker which introduced a Bsp120I site for later manipulations and provided the sequence GGP between the first and second repeat unit. The second repeat was linked to a third repeat by a linker which introduced a BssHII site [again for later manipulation] and resulted in the amino acid sequence GAR. The third repeat was flanked by 2 additional GPP triplets, a GCC triplet and finally GLEGPRG (SEQ ID NO:30). This was a result of including coding sequence that provided for XhoI, SacII and NheI sites. These were included for flexibility of cloning at later stages. The NheI site provides an in frame stop codon.

The synthetic fragment was produced by PCR from primers against COL3A1 in 3 pieces initially. Fragment 1 was EcoRI-NheI-Met-[GAP]2-[REPEAT]1-Bsp120I. The primers for this were 5'-aattccatg-gtgctccaggtgctcc-3' [up] (SEQ ID NO:31)[primer U101] and 5'-ggcc-acctggtggacctggtgg-3' [down] (SEQ ID NO:32)[primer D101]. The second PCR fragment used primers 5'-ggccc-gtgtgtaagggtgacgc-3' [up] (SEQ ID NO:33)[primer U102] and 5'-cgcg-acctggtggacctgg-3' [down] (SEQ ID NO:34)[primer D102]. For the 3rd repeat primer pairs used were 5'-cgcg-gtgtgtaagggtgacgtgg-3' [up] (SEQ ID NO:35)[primer U103] and 5'-acaacctggtggacctggtggacc-tggtggacctgggtgg-3' [down] (SEQ ID NO:36)[primer D103]. The three fragments from the PCR reactions were gel purified and ligated together. The DNA from the ligation mixture was then used as the template for a further round of PCR using primer U101 and a new primer at the 3' end [5'-ctagccccgcggaccctcgagaccaca-acaacctggtgg-3' ] [down] (SEQ ID NO:37)[primer D104]. A band of approximately 500 bp was produced and gel purified, digested with EcoRI-NheI and ligated to pYX141 (Ingenous Cat. No MBV-025-10) [LEU2-CEN-p786] also digested with EcoRI-NheI before being transformed into *E.coli*. Transformants were screened by PCR using primers for the second fragment and DNA from positive colonies were miniprepmed and screened by enzyme digestion with EcoRI-NheI for the presence of an insert of approximate 500 bp. This storage vector



was designated pYX-SYN-C3-1. The EcoRI-NheI fragment was transferred to pYX243 [2u-LEU2-pGAL] (Ingenious Cat. No MBV-035-10) to give pYX-SYN-C3-2 and this plasmid was introduced into a yeast host cell including neucleotide sequence for the carrying the P4H  $\alpha$  and  $\beta$  subunits [either pYEura3.2.12 $\beta$ #39 $\alpha$ #5 or pYAC $\alpha\beta$ ]. Expression following galactose induction was determined by using a MAb 2G8/B1 (Werkmeister & Ramshaw, 1991) which recognises the sequence GLAGAPGLR (SEQ ID NO:38). An EcoRI-SacII fragment from pYX-SYN-C3-2 was also introduced into the EcoRI-SacII of YEplFlag to produce YEplFlag-SYN-C3 and this too was introduced into a yeast host cell expressing P4H on induction by galactose. A product of approximately 18 kDa [the expected size of SYN-C3] was detected in yeast induced with galactose by Western blotting.

The nucleotide sequence for SYN-C3 is provided at Figure 8 together with the amino acid sequence of the encoded product.

#### **Example 8: The use of yeast other than *Saccharomyces cerevisiae*.**

The GAL1-10 promoter is functional in *Kluyveromyces* whilst the *ADH2* promoter is constitutively expressed in *S. pombe*. By shifting the expression cassettes to appropriate vectors, other yeast hosts can be used. *K. lactis* for instance has been shown in some instances to display less proteolytic activity for recombinant products. Alternatively, *P. pastoris* could be used for multiple integration of the expression cassette for  $\alpha$   $\beta$  into the chromosome.

For expression in *P. pastoris*, the nucleotide sequence described in the previous example encoding the synthetic triple helical protein [SYN-C3] was inserted into the *P. pastoris* vector pPIC9 (Invitrogen, Cat. No. K1710-01) at the EcoRI-NotI sites [pPIC-SYN-C3]. Following digestion with either BglII or Sall, the plasmid was introduced into *P. pastoris* where it was integrated at either the AOX1 or HIS4 sites for BglII or Sall respectively. The nucleotide sequences encoding the P4H  $\alpha$  and  $\beta$  subunits were also introduced into *P. pastoris* using the EcoRI site of pHIL-D2 (Invitrogen, Cat. No. K1710-01) for the  $\beta$  subunit and integration at HIS4 and the BamHI site of pHIL-S1 for the  $\alpha$  subunit and subsequent integration HIS4. All three expression cassettes were under the control of the AOX1 promoter and induced by methanol.

**Example 9: Enhanced expression of proly-4-hydroxylase  $\alpha$  and  $\beta$  subunits from the GAL1-10 promoter by use of yeast with different backgrounds for control of galactose induced expression.**

The plasmid pYEUra3.2.12 $\beta$ #39 $\alpha$ #5 [encoding the  $\alpha$  and  $\beta$  subunits of P4H under the control of the GAL1-10 bidirectional promoter] can be introduced into a yeast host cell with the following genotype : a or  $\alpha$ , ura3 trp1 egd1 btt1. In these cells, the absence of the products for the EGD1 and BTT1 genes results in higher levels of galactose induced expression from GAL4 dependent promoters such as GAL2, GAL4, GAL7, GAL1-10, MEL1 (Hu & Ronne, 1994).

Another mechanism for enhanced expression is the use of a yeast host cell carrying multiple copies of the GAL4 (Johnston & Hopper, 1982) positive transcriptional activator under its own controlled induction by galactose. This leads to enhanced expression as there is no limit to the availability of the transcriptional activator for the GAL1-10 promoter. Similarly, the yeast host cell could contain multiple copies of the SGE1 gene (Amakasu *et al.*, 1993) which also leads to enhanced transcription from galactose induced promoters.

Various combinations of these backgrounds could also be utilised; that is egd1 btt1 SGE1<sup>mc</sup> or egd1 btt1 GAL4<sup>mc</sup> or egd1 btt1 SGE1<sup>mc</sup> GAL4<sup>mc</sup> [where mc represents multiple copies].

**Example 10: Expression of collagen from promoters other than ADH2.**

The collagen encoding nucleotide sequence in YEpFlag COL 1.6kb can be excised as a NheI or HindIII- XbaI or NruI fragment for insertion into other fusion vectors under the control of other promoters. Alternatively, the pADH2- $\alpha$  signal-A-proregion-Flag collagen cassette can be excised as a NaeI or SacI - BglII or XbaI or SpeI or SnaBI or NotI, for example, and introduced into an appropriate vector such as YEplac181 (Gietz & Sugino, 1988) or pMH158 (Heuterspreute *et al.*, 1985) for expression in different copy numbers and host backgrounds or into vectors with CEN sequences. Alternatively, CEN sequences can be introduced into the YEpFlag vector itself. The cassette can also be removed without the ADH2 promoter using NruI and introduced into an appropriate vector behind an appropriate promoter.

Collagen encoding nucleotide sequences can be expressed using the CUP1 promoter in vectors such as pYELC5 (Macreadie *et al.*, 1989) as an alternative to the ADH2 promoter. This promoter is induced by addition of copper (i.e. copper sulfate) and may have the advantage of an increased reducing environment and enhancement of P4H activity during co-expression. A second promoter that can be used is the TIP1

promoter which is induced by cold shock. Here the stability of the expressed collagen may be enhanced without the need for hydroxylation by inducing expression by shifting growing yeast from 30°C to 18°C.

#### 5 **Example 11:**

Two single strand synthetic oligonucleotide sequences encoding complimentary forward and reverse strands with complimentary, non-palindromic 5' overhanging sequences were annealed to produce a double-stranded oligonucleotide molecule with the following nucleotide sequence:

10

5' -AGA TCC GGT GGT AAG GGT GAC GCT GGT GCT CCA GGT GAA AGA  
CCA CCA TTC CCA CTG CGA CCA CGA GGT CCA CTT TCT

15

GGT CCA CCA GGT TTG GCT GGT GCT CCA GGT TTG AGA GGT GGT GCT  
CCA GGT GGT CCA AAC CGA CCA CGA GGT CCA AAC TCT CCA CCA CGA

GGT CCA CCA GGT CCA GAA GGT GGT AAG GGT GCT GCT GGT CCA CCA  
CCA GGT GGT CCA GGT CTT CCA CCA TTC CCA CGA CGA CCA GGT GGT

20

GGT CCA CCA GGT (SEQ ID NO:44)  
CCA GGT GGT CCA TCT AGG-5' (SEQ ID NO:45)

which gives rise to the repeat unit sequence;

25

RSGGKGDAGAPGERGPPGLAGAPGLRGGAGPPGPEGGKGAAGPPGPPG (SEQ ID NO:46).

30

Following annealing, the oligonucleotides were then subjected to ligation in the presence of T4 DNA ligase and fractionated by size. Either of the following procedures were followed to clone the various repeat units into YepFlag1 which had been pre-digested with BglII and BamHI.

In the first approach, the following two linkers were added;

35

5' -GATCCGGT (SEQ ID NO:47)  
GCCATCTAGG-5' (SEQ ID NO:48) to anneal at the forward end  
and

5' -AGATCCGGTA (SEQ ID NO:49)

GCCATCTAG-5' (SEQ ID NO:50)

to anneal at reverse strand end.

5

Flag

Linker

**DYKDDDDDKEFLEPGRS** (GRSGGKGDAGAPGERGPPGLAGAPGLRGGAGPPGPEGGKG

10

Linker

AAGPPGPP)nGRSGPVDPR (SEQ ID NO:51).

15

*Flag*

Linker

Linker

**DYKDDDDDKFLEPRS** (GRSGGKGDAGAPGERGPPGLAGAPGLRGGAGPPGPEGGKGAAGPPGPP) **\_GRS**

**IDGSGPVDPR** (SEQ ID NO:52)

20

Expression was as described in Example 7.

25

The method according to the invention provides for the stable expression of triple helical proteins from yeast host cells. Synthetic products may show enhanced or novel functions (e.g. inclusion of RGD and/or YIGSR sequences from fibronectin and laminin, and a fusion of a collagen sequence to a platelet derived growth factor (PDGF) will provide a protein product useful in wound healing and fibrosis). The products may be used in a wide range of applications including bioimplant production, soft and hard tissue augmentation, wound/burn dressings, sphincter augmentation for urinary incontinence and gastric reflux, periodontal disease, vascular grafts, drug delivery systems, cell delivery systems for natural factors and as conduits in nerve regeneration.

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It will be appreciated by persons skilled in the art that numerous variations and/or modifications may be made to the invention as shown in the specific embodiments without departing from the spirit or scope of the invention as broadly described. The present embodiments are, therefore, to be considered in all respects as  
20 illustrative and not restrictive.